Kathrin Schmidt<sup>a)</sup>, Ronald Gust<sup>a)</sup>, and Manfred Jung\* b)

a) Institut für Pharmazie der Freien Universität Berlin, Abteilung Pharmazeutische Chemie, Königin-Luise-Str. 2 + 4, 14195 Berlin, Germany

b) Institut für Pharmazeutische Chemie, Westfällische Wilhelms-Universität Münster, Hittorfstr. 58-62, 48149 Münster, Germany

Key Words: Histone deacetylase; trichostatin A: cancer: MCF-7

#### Summary

Inhibition of histone deacetylase are attracting increasing interest due to their influence on transcription, differentiation, and proposition. We have investigated two synthetic inhibitors 3 and 4 of the histone deacetylase and the natural protect inhibitor includes and the natural protect inhibitor includes and for their ability to suppress the growth of MCF-7 breast camer cells and here present complete and improved synthetic procedures. The compounds show a dose dependent inhibition of growth in activities in the low micromologia and nanonolar range. Trichostatin shows extectical effects at 100 mM and still has settivity comparable to edgalatin (0.5 LMM) at 10 MM. Whereas the synthetic inhibitor 3 has cytocidal activity at 10 JM compound 4 shows a maximum of 40% growth suppression at that exceptantians that exceptantians and that exceptantians and that exceptantians and that exceptantians after acceptantians.

#### Introduction

Reversible acetylation of e-amine groups of highly conserved lysine residues in the N-terminal tails of histones has a modulating impact on chromatin structure. This affects key nuclear processes such as DNA replication, transcription, differentiation, and apoptosis "In his modification is established and maintained by histone acetyltransferrases and histone deacetylases, enzymes which have been identified as homologo of regulators of transcription and nucleolar phosphoproteins E-"uncreasing evidence shows that transcription factors involved in the regulation of proliferation and differentiation exert their function by recruitment of histone acetyltransferases or deacetylases [2,3,5,]. In several cases aberrant chromatin acetylation could be linked to malignant diseases [5,7-9].

In the course of our studies to synthesize simple inhibitors of histone deacetylase and the exploration of their potential for cancer therapy and chemoprevention we have found promising lead substances <sup>[10]</sup>. Among them were compounds 3 and 4 constructed from structural elements of the established natural product inhibitors trapoxin B (1) and trichostatin A (2) (Scheme I) <sup>[10]</sup>

Whereas the impact of inhibitors of histone deacetylase on leukaemic cells has been studied at length comparatively littleis known about their influence on cell lines derived from solid tumors. Inst recently it has been shown that the auticuscer drug cisplatin is crossificking molear proteins to the District drug cisplatin is crossificking molear proteins to the District MCF-7 breast cancer cells and it was suggested that its clinical activity is linked partly to that effect. Among those nuclear proteins were the estrogen receptor and histone deacetylase HDAC-1<sup>[12]</sup>. So we studied the growth inhibition by reference histone deacetylase inhibitor trichostatin A (2) and of our synthetic lead structures 3 and 4 on that human breast cancer cell line.

### Results and Discussion

## Chemistry

As described previously 3 and 4 are accessible in four synthetic steps using standard peptide coupling methodology.

Scheme 1. Inhibitors of histone deacetylase.

Scheme 2. Synthesis of Inhibitors 3 and 4.

As the target compounds are very polar we chose a strategy where no chromotography is required after the last synthetic step. Thus a benzyl-protected hydroxylamine is employed that allows for chromotographic purification of the protected hydroxamate. After hydrogenation the compounds can be purified by precipitation from methanolic solutions. The use of BOP-C1 in all coupling steps in the synthesis of 3 led to increased yields compared to the water-soluble carbodimide EDC. For 4 we had afteraby proven that the use of BOP-C1 is crucial in the first coupling steps a mixed analydide or carbodimide methods failed due to the inactivating effect of the para-dimethylamino group "IO" (Sechem 2.).

#### Biological Activities

The impact of the different inhibitors of histone deacetylase on MCF-7 cells was studied in our standard microtiter assay using crystal violet staining to measure total biomass relative to control over a prolonged period of exposure to the antineoplastic agent [13] Cisplatin was used as a reference compound (Figure 1).

Trichostatin A (2) and inhibitor 3 are able to reach systocidal effects at 10 on M(2) and 10 µM (3), respectively. In contrast, compound 4 did not exceed 40% suppression of cell growth even at 10 µM. Even at 10 µM, 2 still has an activity comparable to cisplaint a 300 nM. This difference between 2 and our inhibitions suppression (250 acts 2 nM, 3 soo) nM, 4: 100 nM) <sup>110</sup>. After approximately 60 h there was an increase in cell growth in some of the experiments (2 at 10 nM, 3 at 10 µM, 4 at 10 µM) which may reflect hydrolysis of the hydroxamates to the inactive carboxylic acids.

Although the dose response curves are rather steep it is encouraging that inhibitors of thisone deacetybase can suppress the growth of breast cancer cells and even show evicial effects in some cases. It has been shown recently that this enzyme can indeed be inhibited safely in humans. The depsipeptide FR-901228 which was chosen for clinical trials due to its good antitumor activity was discovered to be an inhibitor of histone deacetylase [13]. First results from clinical brail plane 1 studies showed encouragingly low toxicity [16].

Moreover, a patient with multi-resistent acute promyelocytic leukaemia was found to respond again to retinoid treatment by combination with the unspecific histone deacetylase inhibitor phenybluyrate. Six months after teatment RT-PCR did not show any mRNA of the oncogenically transformed retinoid receptors that recruit histone deacetylase is superses transcription, thus indicating absence of residual disease. Again no severe side effects were observed <sup>117</sup>. Thus this first report of the growth inhibition of MCF-7 breast cancer cells by potent inhibitors of histone deacetylase and so the mountain of the mountain of the second of the provident of

#### Acknowledgements

M. Jung wishes to thank Prof. Unterhalt, Münster for his support and H. Deguara for technical sasistance. Financial support by the Fonds der Chemischen Industrie, the DPHo-Sithung flut en wissenschaftlichen Nach-wuchs im Stifterverbund für die Deutsche Wissenschaft, the Förderegsenlischaft der Westfallischen Wilhelms-Universität and the Deutsche Forschungsgemeinschaft is gratefully acknowledged. We thank ASTA Medica AG, Frankfurf, for gifts of S-miniocappic acid and 2.

#### Experimental Part

Melting point: uncorrected.— Elemental analysis: Perkin Elmer.— IR: Shimadru 470.— H-NMR: Varian Gemini 200 (200 MHz).— <sup>13</sup>C-NMR: Varian Gemini 200 (2029 MHz).— MS: Varian MAT 44S, Fimigan MAT 312.— Flash chromatography (FC): silke gel 60, 230-400 mesh (Merck).— Dichloromethane was dried over molecular sieves (3.8).

## Chemistry

#### General Procedure for Amide Formation

To a solution or suspension of the acid in dry CH<sub>2</sub>Cl<sub>2</sub> (5 ml/mmol) was added one equivalent of triethylamine (TEA) under nitrogen and the mixture was stirred for 10 min. Then 1.1 equivalents of BOP-Cl, one equivalent of the amine or hydroxylamine hydrochloride, and another three equivalents of

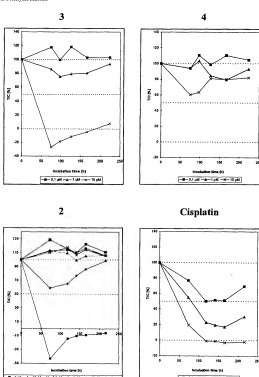


Figure 1. Growth inhibition of MCF-7 breast cancer cells by inhibitors of histone deacetylase.

General method for amide formation: 199 g. (10 mmol) monomethyl substrata, 1.39 m. (10, g. 10 mmol) Tax. P.2. 30 g. (11 mmol) BOPC, 1.60 g. (10 mmol) O-Benzyllhydroxyllamine bydrochloride, 4.17 m. I. (3.0 g. g. 10 mmol) Tax. Read on wasted with 2.0 mmol Tax. Read on wasted with 2.0 mmol Tax. Read on wasted with 2.0 mmol Tax. Read on the control of t

#### N-Benzyloxymonosuberoylamide (5)

2.77 g (10 mmo) of methy) M-benzylo-synberoy/smide were dissolved in Om. of THE and I om. (20 mmo) of a Ht LiGH Solvinie were added. The missue was stirred for 4 h. at room temperature. 10 m. of 0.5 M NaOH and 50 mt. of ethyl seather were added. The sequence plates was saddified and the control of the con

### N-Benzyloxy-N-[1-(S)-methoxycarbonyl-2-phonylethyl]suberoylbisamide

General method for amide formation; 1.11 g (4 mmol) 5, 0.5 mL (0.41 g, 4 mmol) TEA, 0.86 g (4 mmol) L-phenylalanine methyl ester hydrochloride, 1.12 g (4.4 mmol) BOP-Cl, 1.68 mL (1.21 g, 12 mmol) TEA. The organic phase was also washed with 2 M HCl. The crude was chromatographed using ethyl acetate/hexane (5:1). Yield 1.25 g (71%) colorless crystals.- Mp 86 °C .- IR (KBr): v = 3050s, 1730s, 1640s.- MS (70 eV); m/r (%) = 440 (3) [M\*1, 274 (14), 162 (100),- 1H NMR (CDCh); δ = 8.79 (bs. 1 H. NHOCHs). 7.37 (s, 5 H, OCH<sub>2</sub>Ph), 7.33-7.22 (m, 3 H, Phe-H), 7.11-7.06 (m, 2 H, Phe-H), 6.02 (d, 3J = 7.53 Hz, 1 H, Phe-NH), 4.89-4.81 (m, 3 H, OCH2 and CHCOOMe), 3.71 (s, 3 H, OMe), 3.19-3.02 (m, 2 H, CHCH2Ph), 2.17-2.09 (m, 2 H, CH<sub>2</sub>), 2.02 (bs, 2 H, CH<sub>2</sub>), 1.63-1.49 (m, 4 H, CH<sub>2</sub>), 1.27-1.24 (m, <sup>13</sup>C NMR (CDCl<sub>3</sub>): δ = 172.70 (COOMe), 172.31 (CONH), 4 H, CH2) .-135.99 (Phe-1), 135.00 (OCH<sub>2</sub>C), 129.22, 129.18, 128.70, 128.60 and 127.14 (Ar-CH), 78.33 (OCH2), 53.06 (OMe), 52.27 (CHCOOMe), 37.94, 36.18, 32.70, 28.48, 25.20 and 24.82 (CH2). Anal. (C25H32N2O5) 440.59: C, H, N.

### N-Hydroxy-N-[1-(S)-methoxycorbonyl-2-phenylethyl]suberoylbisomick (3)

1.00 g. (2.27 mmol). N-bernyloxyN-(1-(35)-methoxycarbosy)-2-bepryleythyl-shroyblismide was disovide in 15 m. of methanol and 100 mg 10% pallodium on churcoal were added. The mixture was treatment with hydrogen under atmospheric pressure for 6 has also hasquestry filtered. The product was evaporated, redissolved in 5 mL of methanol, and precipitate with indirely inter. Yield 310 m. (94%) colorises regulate. Mp 105 °C-, 18 (XB): y = 320m. 1790., 1809.—185 (70 eV); not (9): −3.00 (19): 1.00 (19): 1.00 (19). 1.00 (19): 1.

36.63, 34.89, 32.21, 28.30, 28.11 and 24.97 (CH<sub>2</sub>, one resonance is obscured).- Anal. (C<sub>14</sub>H<sub>26</sub>N<sub>2</sub>O<sub>3</sub>) 350.46; C. H. N.

### Methyl 6-(4-Dimethylaminobenzoylamino)caproote

General method for amide formation: 1.5.5 g (10 manol) 4-dimethylamino bruncia caid, 1.3 pd. (10 manol) method formation: 1.5.5 g (10 manol) method formation: 1.5.2 g (10 manol) method formation: 1.5.2 g (10 manol) method formation for

#### 6-(4-Dimethylaminobenzoylamino)caproic Acid (6)

1.17 g (4 mmol) of methyl 6-(4-dimethylaminobenzoylamino)caproste were dissolved in 10 mL of THF and 4 mL (8 mmol) of a 1 M LiOH soluti were added. The mixture was stirred for 4 h at room temperature, 10 mL of 0.5 M NaOH and 50 mL of ethyl acetate were added. The aqueous phase was treated with 2 M HCl to pH 6-7 and some of the scid precipitated. The mother liquor was evaporated to dryness and dried over P2O5 in a desiccator overnight. The resulting solid was sufficiently pure for the next coupling step. The spectral data was taken form the precipitated pure acid.- Mp 165 °C .-IR (KBr): v = 3380s, 2910s, 1700s, 1600s,- MS (70 eV); m/z (%) = 278 (36)  $[M^{+}]$ , 219 (19), 148 (100).— H NMR (CD<sub>3</sub>OD):  $\delta = 7.71-7.66$  (m, 2 H, 2'-H, 6'-H,), 6.75-6.70 (m, 2 H, 3'-H, 5'-H), 3.37-3.28 (m, 2 H, CONHCH2), 3.01 (DMSO-d6): δ = 174.32 (COOH), 165.97 (CONH), 151.94 (Me<sub>2</sub>NC), 128.34 (C-3', C-5'), 121.48 (CCONH), 110.71 (C-2', C-6'), 39.65 (NMe<sub>2</sub>), 38.81 (CONHCH2), 33.59 (CH2COOMe), 29.02, 26.02, 24.23 (CH2).- Anal. (C15H18N2O3) 278.32: C, H, N.

### N-Benzyloxy-6-(4-dimethylaminobenzoylamino)capramide

General method for analysis formation; combined and crude asportification productd, 6.5 mil. (ed. 8, ed. mon) TEAS, 0.6 § 6 mmol) Developidyncty-lamine hydrockhloride, 1.12 g (4.4 mmol) BCP-Cl, 1.68 mil. (1.21 g. 12 mmol), TEA. The crude was chromotographed using ently aceste with 1% distributions in 18 mil. (ed. 18 mil. 18

# N-Hydroxy-6-(4-dimethylaminobenzoylamino)capramide (4)

333 mg (1 mmol) of //bemploxy-6-(4-dimethylminobemoylamino) capmanide were dissolved in 15 mL of relational and 40 mg (16% palladium on charcoal were added. The initiate was treated with hydrogen under amospheric pressure for 6 has dwa subleagenshy filtered. The product was evaporated, redissolved in 5 mL of methanol, and precipitated with diethyl prior 10 mL of 10 mL o

#### Biological Methods

#### Cell Culture

The human MCF-2 breast cancer cell line was obtained from the American Dye Culture Collection (ATCC, Rockstill, Md. 128A), Cell line backing, and quality control were performed according to the send stock concept verieved by Hay, 10<sup>3</sup>. The MCF-7 cell were maintained as a monelayer culture at 37° C in a humidified 95% at i, 55° CO<sub>2</sub> atmosphere in 1-75 flasks using L-plustames constaining Eaglet, MMS supplemented with BCS (100mLIA) as growth medium. The cell line was weekly passaged after treatment with specific DTA.

### In Vitro Chemosensitivity Assay

The no vicro testing of the inhibition of histone descrylate for antitumor activity was carried on MCF-7 cells recording to a previously published micrositer assay <sup>10</sup>. Bretly, 100 aL. of a cell suspension at 500 cellium-Leutem medium were planed into an who of a 68-west minorities phea and cauties medium were planed into an elven of a 68-west minorities phea and substance was added in three concentrations. After the proper incubations in substance was added in three concentrations. After the proper incubations in the medium was removed, the cells were fixed with a plantial disable solution and stood of 4° C. Cell blomas was determined by a crystal violate that the contract of the contract of

$$T/C_{corr}$$
 [%] =  $[(T - C_0) / (C - C_0)] \times 100$ 

where T (test) and C (control) are the optical densities at 590 nm of the crystal violet extract of the cells in the wells  $(c. the chromatin-bound crystal violet extracted with ethanol 70%), and <math>C_0$  is the optical density of the cell extract immediately before treatment. Cytocidal effect:

$$T_{corr} [\%] = [(T - C_0) / C_0] \times 100$$

For the automatic estimation of the optical density of the crystal violet extract in the wells the Microplate Photometer Labsystem Multisean® Plus was used.

### References

- [1] P. Loidl, Chromosoma 1994, 103, 441-449,
- [2] E. Pennisi, Science, 1997, 275, 155-157.

- [3] M. J. Pazin, J. T. Kadanoga, Cell 1997, 86, 325-328.
- [4] A. Lusser, G. Brosch, A. Loidl, H. Haas, P. Loidl, Science 1997, 277, 88-91.
- [5] K. Struhl, Genes Dev. 1998, 12, 599-606.
- [6] R. J. Liu, L. Nagy, S. Inoue, W. Shao, V. H. Miller, R. M. Evans, Nature 1998, 391, 811–814.
- [7] F. Grignani, S. De Matteis, C. Nervi, L. Tomassoni, V. Gelmetti, M. Cioce, M. Fanelli, M. Ruthardt, F. F. Ferrara, I. Zamir, C. Seiser, F. Grignani, M. A. Lazar, S. Minucci, P. G. Pelicci, *Nature* 1998, 391, 815–818
- [8] J. Borrow, V. P. Stanton Jr., J. M. Andresen, R. Becher, F. G. Behm, R. S. Chaganti, C. I. Civin, C. Disteche, I. Dube, A. M. Frischauf, D. Horsman, F. Mitelman, S. Volinia, A. E. Watmore, D.E. Housman, Nature Genetics 1996, 14, 33-41.
- [9] H. Siddique, J.-P. Zou, V.N. Rao, E.S.P. Reddy, Oncogene 1998, 16, 2283–2285.
- [10] M. Jung, K. Hoffmann, G. Brosch, P. Loidl, Bioorg. Med. Chem. Lett. 1997, 7, 1655-1658.
- [11] M. Yoshida, S. Horinuchi, T. Beppu, Bioassays 1995, 17, 423–430.
- [12] S. K. Samuel, V. A. Spencer, L. Bajno, J. M. Sun, L. T. Holth, S. Oesterreich, J. R. Davie, Cancer Res. 1998, 58, 3004–3008
- [13] G. Bernhardt, H. Reile, H. Bimböck, T. Spruß, H. Schönenberger, J. Cancer Res, Clin. Oncol. 1992, 118, 35–43.
- [14] K. Hoffmann, G. Brosch, P. Loidl, M. Jung, Pharmazie, submitted.
- [15] H. Nakayima, Y. B. Kim, H. Terano, M. Yoshida, S. Horinouchi, Experimental Cell Res. 1998, 241, 126-133.
- [16] J. L. Marshall, W. L. Dehut, N. Rizvi, I. W. Wainer, C. Chassaing, M. Figiulera, M. J. Hawkins, 10<sup>th</sup> NCI-EORTC-Symposium on new drugs in cancer therapy, 16–19, 06, 1998, Amsterdam.
- [17] R. P. Warrell Jr., L-Z. He, V. Richon, E. Calleja, P. P. Pandolfi, J. Natl. Cancer Inst, 1999, 90, 1621-1625.
- [18] J. Taunton, C. A. Hassig, S. L. Schreiber, Science 1996, 272, 408-411.
- [19] B. Nare, J. J. Allocco, R. Kuningas, S. Galuska, R. W. Myers, M. A. Bednarek, D. M. Schmatz, Anal. Biochem. 1999, 267, 390-396.
- [20] K. Hoffmann, G. Brosch, P. Loidl, M. Jung, Nucleic Acids Res. 1999, 27, 2057–2058.
- [21] R. J. Hay, Anal. Biochem. 1988, 171, 225-237.
- [22] H. Reile, H. Birnböck, G. Bernhardt, T. Spruß, H. Schönenberger, Anal. Biochem. 1990, 187, 262-267.

Received: May 6, 1999 [FP394]